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Process to prepare a lubricating base oil

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## PROCESS TO PREPARE A LUBRICATING BASE OIL

The invention is directed to a process to prepare a base oil having a viscosity index of above 80 and a saturates content of above 90 wt% from a crude derived feedstock by means of a process comprising a hydrocracking step and a catalytic dewaxing step.

EP-A-0909304 illustrates a process wherein a base oil having a viscosity index (VI) of 95 is prepared from a vacuum distillate boiling between 418 (5 wt% recovery) and 564 °C (95 wt% recovery) by subjecting the feed to a hydrocracking step using a catalyst based on Nickel and Molybdenum. The high boiling part of the hydrocracker effluent was subsequently dewaxed using a ZSM-5 based dewaxing catalyst and hydrofinished using a platinum/palladium based catalyst. The yield to base oil was 62 wt%.

WO-A-0250213 describes a process to prepare a base oil from the high boiling fraction of a fuels hydrocracker process. In this process the high boiling fraction is separated into different distillate fractions which are in turn subjected to a catalytic dewaxing step and a hydrofinishing step.

US-A-5525209 describes a fuels hydrocracker process wherein the bottoms fraction in which bottoms fraction may potentially yield a base oil having a desired high viscosity index value. It is shown in this publication that the viscosity index of the base oil will increase at higher conversion in the hydrocracker step.

According to general textbooks on base oil manufacturing hydrocracking will reduce the viscosity of the feedstock, remove most of the nitrogen, oxygen and

5           sulphur present in the base oil feedstock and convert the undesirable low VI materials such as polynuclear aromatics and polynuclear naphthenes to higher VI materials such as mononuclear aromatics, mononuclear naphthenes and iso-paraffins (Chapter 6 and especially page 122 of Lubricant Base Oil and Wax Processing, Avilino Sequeira, Jr, Marcel Dekker Inc, New York, 1994, ISBN 0-8247-9256-4).

10          A disadvantage of the above processes is that not all crude derived feedstocks are suitable for preparing a base oil having the desired VI. It may also be possible that a crude derived feed is suitable to meet the VI requirements of some but not all of the desired viscosity grades. This could for example be due to the fact that  
15          the content of polynuclear aromatics and naphthenics in the relevant feed or feed fraction are too high. It may sometimes be possible to meet the VI requirements by increasing the hydrocracker conversion as explained above. However such a higher conversion will  
20          significantly lower the final base oil yield and may even make it impossible to prepare the heavier grades.

The object of the present invention is to provide a process, which overcomes the above-described disadvantages of the prior art.

25          This object is achieved with the following process. Process to prepare a base oil having a viscosity index of above 80 and a saturates content of above 90 wt% from a crude derived feedstock by

30          (a) Contacting a crude derived feedstock in the presence of hydrogen with a catalyst comprising at least one Group VIB metal component and at least one non-noble Group VIII metal component supported on a refractory oxide carrier;

35          (b) Adding to the effluent of step (a) or part of the ~~effluent~~ of step (a) a Fischer-Tropsch derived fraction

boiling at least partly in the base oil range in an amount effective to achieve the target viscosity index of the final base oil; and

(c) Dewaxing the mixture as obtained in step (b).

5       Applicants found that the use of Fischer-Tropsch fraction in the process according the invention greatly improves the flexibility of the process. Feedstocks derived from crudes which normally did not yield a base oil having the desired VI could now be used and/or the yield of base oil as calculated on the petroleum derived feedstock could be improved.

10      The petroleum derived feedstock as used in step (a) may be a vacuum distillate fraction as obtained from the residue of the atmospheric distillation of a crude 15     petroleum feed. Such a fraction may be a vacuum gas oil or heavier fractions. The residue of the vacuum distillation itself may also be used. Suitably a vacuum residue is used which has been de-asphalting. Other possible feeds are for example the cycle oils as obtained 20     in a fluid catalytic cracking process. Mixtures of the above feeds are of course also possible. If heavy base oil grades are preferred a feed is used wherein more than 10 wt%, preferably more than 20 wt% and most preferably more than 30 wt% of the compounds present in said feed 25     boil above 470 °C. Suitably less than 60 wt% of the compounds present in the feed boil above 470 °C.

The feed to step (a) will typically have a low VI value of below 60 due to the presence of polynuclear aromatics and naphthenics. The VI of the feed as here defined is the VI of a solvent dewaxed sample having a pour point of -18 °C.

30      Step (a) may be performed according to well known hydrocracking processes. These processes may be both hydrocracking processes known to make primarily middle distillates and base oil hydrocracking processes. The 35

conversion in step (a), expressed in the weight percentage of the fraction in the feed which boils above 370 °C which is converted to products boiling below 370 °C, in step (a) may thus range from values typical for base oil hydrocrackers and to values typical for fuels hydrocrackers. Such conversions may thus be between 20 and 80 wt%. The degree of conversion will depend on the feedstock quality as explained above and the availability of the Fischer-Tropsch derived blending fraction. A skilled person will be able to optimise the conversion given these parameters.

Step (a) may in addition also comprise a hydrotreating step performed prior to the actual hydrocracking step. In the hydrotreating step nitrogen and sulphur are removed and aromatics are saturated to naphthenes. The reduction in sulphur and nitrogen is preferably such that the feed to step (c) is below 100 ppmw sulphur and more preferably below 50 ppm sulphur and more preferably below 10 ppmw nitrogen.

It has been found that in the process according to the present invention a base oil may be prepared having the desired VI wherein the conversion in the hydrotreating step is relatively low. This is especially advantageous when also more heavier grades are desired. The conversion is preferably below 40 and more preferably below 30 wt%. The preliminary hydrotreating step is typically performed using catalyst comprising a metal hydrogenation component, suitably a combination of a Group VIB and a non-noble Group VIII metal, for example cobalt-molybdenum, nickel-molybdenum, on a porous support, for example silica-alumina or alumina. The hydrotreating catalysts suitably contains no zeolite material or a very low content of less than 1 wt%. Examples of suitable hydrotreating catalysts are the commercial RCM 106, ICR 120 of Chevron Research and

Technology Co.; 244, 411, DN-120, DN-180, DN-190 and DN-200, DN-3110, DN-3100 and DN-3120 of Criterion Catalyst Co.; TK-555 and TK-565 of Haldor Topsoe A/S; HC-k, HC-P, HC-R and HC-T of UOP; KF-742, KF-752, KF-846, KF-848 STARS and KF-849 of AKZO Nobel/Nippon Ketjen; and HR-438/448 of Procatalyse SA.

The hydrotreating step is suitably performed at the following conditions: temperature of at least 300 °C, preferably from 350 to 450 °C and even more preferably from 370 to 430 °C. Operating pressure may range from 10 to 250 bar, but preferably is at least 80 bar, more preferably at least 110 bar. In a particularly advantageous embodiment the operating pressure is in the range of from 110 to 170 bar. The weight hourly space velocity (WHSV) may range from 0.1 to 10 kg of oil per litre of catalyst per hour (kg/l.h) and suitably is in the range from 0.2 to 5 kg/l.h.

The hydrocracking step may be any hydrocracking process using well known hydrocracking catalysts or variations of such catalysts having a hydrogenation/dehydrogenation function on a suitable support. Such a function is preferably a Group VIII/Group VIB metal combination, for example nickel-molybdenum and nickel-tungsten. The support is preferably a porous support, for example silica-alumina and alumina. The catalyst may also comprise an, optionally partly dealuminated, large pore size zeolite. Examples of suitable zeolites are zeolite X, Y, ZSM-3, ZSM-18, ZSM-20 and zeolite beta of which partly dealuminated zeolite Y is most preferred. Examples of suitable hydrocracking catalysts are the commercial ICR 220 and ICR 142 of Chevron Research and Technology Co; Z-763, Z-863, Z-753, Z-703, Z-803, Z-733, Z-723, Z-673, Z-603 and Z-623 of Zeolist International; TK-931 of Haldor Topsoe A/S; DHC-32, DHC-41, HC-24, HC-26, HC-34 and HC-43 of UOP; KC2600/1, KC2602, KC2610, KC2702 and KC2710 of AKZO

Nobel/Nippon Ketjen; and HYC 642 and HYC 652 of Procatalyse SA.

The hydrocracking step is suitably performed at the following conditions: temperature of at least 300 °C, preferably from 340 to 450 °C and even more preferably from 350 to 430 °C. Operating pressure may range from 10 to 250 bar, but preferably is at least 80 bar, more preferably at least 110 bar. In a particularly advantageous embodiment the operating pressure is in the range of from 110 to 170 bar. The weight hourly space velocity (WHSV) may range from 0.1 to 10 kg of oil per litre of catalyst per hour (kg/l.h) and suitably is in the range from 0.2 to 5 kg/l.h.

In step (b) all or part of the effluent of step (a) is mixed with the Fischer-Tropsch derived fraction. Preferably only the fraction of said effluent boiling in the base oil range is used in step (a). Suitably this fraction has an initial boiling point higher than 300 °C and more preferably higher than 340 °C. A maximum value for the initial boiling point will depend on the desired base oil grade one wishes to prepare.

The Fischer-Tropsch fraction may in principle be any fraction which boils in the base oil range and which is isolated from the synthesis product of the Fischer-Tropsch reaction. More preferably a partly or whole hydroisomerized Fischer-Tropsch wax is used. The use of the isomerised product is preferred because a significant part of the normal paraffins as present in a Fischer-Tropsch synthesis product have then been isomerised to the, for base oil manufacture, more desirable iso-paraffins. The Fischer-Tropsch fraction preferably has a boiling range, which corresponds with the petroleum derived fraction as used in step (b).

The Fischer-Tropsch derived fraction may be obtained by well-known processes, for example the so-called commercial Sasol process, the Shell Middle Distillate Process, etc by the non-commercial Exxon process. These and

other processes are for example described in more detail  
in EP-A-776959, EP-A-668342, US-A-4943672, US-A-5059299,  
WO-A-9934917 and WO-A-9920720. The process will generally  
comprise a Fischer-Tropsch synthesis and a hydro-  
isomerisation step as described in these publications.  
Preferably the fraction will comprise of a substantial  
amount of compounds boiling in the base oil range. The  
fraction preferably has a relatively low pour point,  
which is beneficial when the Fischer-Tropsch fraction has  
to be transported from remote locations to the base oil  
process facility. For this reason the Fischer-Tropsch  
fraction has been partly isomerised. An example of a  
suitable fraction is the so-called Shell MDS Waxy  
Raffinate as obtainable from Shell MDS (Malaysia) Sdn Bhd  
or the product as described in WO-A-02070630 or fractions  
of said products. Alternatively n-paraffin waxes as  
obtainable from said processes may also be used having  
preferably a congealing point of between 20 and 80 °C.  
Examples are SX-30, SX-50 and SX-70 as obtainable from  
Shell MDS (Malaysia) Sdn Bhd. If such waxes are used a  
catalytic dewaxing in step (c) is preferred, more  
preferably a dewaxing catalyst is used having a high  
ability to isomerise the normal paraffins. See for  
preferred catalysts below. Of course fractions having  
similar properties as described above as obtainable from  
other processes may also be advantageously used in our  
invention.

The mixture as obtained in step (b) will suitably  
have a viscosity corresponding to the desired viscosity  
of the base oil product. Preferably the kinematic  
viscosity at 100 °C of the mixture is between 3 and  
10 cSt. The content of Fischer-Tropsch derived fraction  
in the mixture is preferably higher than 5 wt%, more  
preferably higher than 10 wt% and preferably lower than

50 wt% and more preferably below 30 wt% and even more  
preferably below 25 wt%.

With the dewaxing in step (c) is understood every  
process wherein the pour point of the base oil is reduced  
5 by more than 10 °C, preferably more than 20 °C, more  
preferably more than 25 °C. The dewaxing can be performed  
by means of a so-called solvent dewaxing process or by  
means of a catalytic dewaxing process. Solvent dewaxing  
is well known to those skilled in the art and involves  
10 admixture of one or more solvents and/or wax  
precipitating agents with the base oil precursor fraction  
and cooling the mixture to a temperature in the range of  
from -10 °C to -40 °C, preferably in the range of from -  
20 °C to -35 °C, to separate the wax from the oil. The  
15 oil containing the wax is usually filtered through a  
filter cloth which can be made of textile fibres, such as  
cotton; porous metal cloth; or cloth made of synthetic  
materials. Examples of solvents which may be employed in  
the solvent dewaxing process are C<sub>3</sub>-C<sub>6</sub> ketones (e.g.  
20 methyl ethyl ketone, methyl isobutyl ketone and mixtures  
thereof), C<sub>6</sub>-C<sub>10</sub> aromatic hydrocarbons (e.g. toluene),  
mixtures of ketones and aromatics (e.g. methyl ethyl  
ketone and toluene), autorefrigerative solvents such as  
liquefied, normally gaseous C<sub>2</sub>-C<sub>4</sub> hydrocarbons such as  
25 propane, propylene, butane, butylene and mixtures  
thereof. Mixtures of methyl ethyl ketone and toluene or  
methyl ethyl ketone and methyl isobutyl ketone are  
generally preferred. Examples of these and other suitable  
solvent dewaxing processes are described in Lubricant  
30 Base Oil and Wax Processing, Avilino Sequeira, Jr, Marcel  
Dekker Inc., New York, 1994, Chapter 7.

Preferably step (c) is performed by means of a  
catalytic dewaxing process. The catalytic dewaxing  
step (c) can be performed by any process wherein in the  
presence of a catalyst and hydrogen the pour point of the

base oil precursor fraction is reduced as specified above. Suitable dewaxing catalysts are heterogeneous catalysts comprising a molecular sieve and optionally in combination with a metal having a hydrogenation function, such as the Group VIII metals. Molecular sieves, and more suitably intermediate pore size zeolites, have shown a good catalytic ability to reduce the pour point of the distillate base oil precursor fraction under catalytic dewaxing conditions. Preferably the intermediate pore size zeolites have a pore diameter of between 0.35 and 0.8 nm. Suitable intermediate pore size zeolites are mordenite, ZSM-5, ZSM-12, ZSM-22, ZSM-23, SSZ-32, ZSM-35 and ZSM-48. Catalysts having a high ability to isomerise normal paraffins will preferably comprise ZSM-12, ZSM-22, ZSM-23 or SSZ-32. Another preferred group of molecular sieves are the silica-aluminaphosphate (SAPO) materials of which SAPO-11 is most preferred as for example described in US-A-4859311. ZSM-5 may optionally be used in its HZSM-5 form in the absence of any Group VIII metal. The other molecular sieves are preferably used in combination with an added Group VIII metal. Suitable Group VIII metals are nickel, cobalt, platinum and palladium. Examples of possible combinations are Ni/ZSM-5, Pt/ZSM-23, Pd/ZSM-23, Pt/ZSM-48 and Pt/SAPO-11. Further details and examples of suitable molecular sieves and dewaxing conditions are for example described in WO-A-9718278, US-A-5053373, US-A-5252527 and US-A-4574043.

The dewaxing catalyst suitably also comprises a binder. The binder can be a synthetic or naturally occurring (inorganic) substance, for example clay, silica and/or metal oxides. Natural occurring clays are for example of the montmorillonite and kaolin families. The binder is preferably a porous binder material, for example a refractory oxide of which examples are:

alumina, silica-alumina, silica-magnesia, silica-zirconia, silica-thoria, silica-beryllia, silica-titania as well as ternary compositions for example silica-alumina-thoria, silica-alumina-zirconia, silica-alumina-magnesia and silica-magnesia-zirconia. More preferably a low acidity refractory oxide binder material, which is essentially free of alumina, is used. Examples of these binder materials are silica, zirconia, titanium dioxide, germanium dioxide, boria and mixtures of two or more of these of which examples are listed above. The most preferred binder is silica.

A preferred class of dewaxing catalysts comprise intermediate zeolite crystallites as described above and a low acidity refractory oxide binder material which is essentially free of alumina as described above, wherein the surface of the aluminosilicate zeolite crystallites has been modified by subjecting the aluminosilicate zeolite crystallites to a surface dealumination treatment. A preferred dealumination treatment is by contacting an extrudate of the binder and the zeolite with an aqueous solution of a fluorosilicate salt as described in for example US-A-5157191 or WO-A-2000029511. Examples of suitable dewaxing catalysts as described above are silica bound and dealuminated Pt/ZSM-5, silica bound and dealuminated Pt/ZSM-23, silica bound and dealuminated Pt/ZSM-12, silica bound and dealuminated Pt/ZSM-22 as for example described in WO-A-200029511 and EP-B-832171.

Catalytic dewaxing conditions are known in the art and typically involve operating temperatures in the range of from 200 to 500 °C, suitably from 250 to 400 °C, hydrogen pressures in the range of from 10 to 200 bar, preferably from 40 to 170 bar, weight hourly space velocities (WHSV) in the range of from 0.1 to 10 kg of oil per litre of catalyst per hour (kg/l/hr), suitably

from 0.2 to 5 kg/l/hr, more suitably from 0.5 to 3 kg/l/hr and hydrogen to oil ratios in the range of from 100 to 2,000 litres of hydrogen per litre of oil. By varying the temperature between 315 and 375 °C at between 5 40-70 bars, in the catalytic dewaxing step it is possible to prepare base oils having different pour point specifications varying from suitably lower than -60 to -10 °C.

10 If the feed to a catalytic dewaxing step (c) has a relatively high nitrogen content of above 10 ppm a pre-treat step is preferably performed wherein under hydroconversion conditions similar to the dewaxing conditions the feed to step (c) is contacted with a noble metal catalyst. Examples of suitable noble metal catalysts are the palladium/platinum containing catalysts C-624 and C-654 of Criterion Catalyst Company. After such 15 a treatment the nitrogen content is reduced to below 10 ppm that is advantageous for the performance of the dewaxing catalyst downstream said treatment.

20 After performing the pour point reducing treatment lower boiling compounds formed during said treatment are suitably removed, preferably by means of distillation, optionally in combination with an initial flashing step.

25 The effluent of the pour point reducing treatment may suitably be subjected to a hydrogenation treatment step (d). Hydrogenation may be performed on the entire effluent or on specific base oil grades after the above-described fractionation. This may be required in order to increase the content of saturate compounds to values 30 above 90 wt% more preferably above 95 wt%. Such a hydrogenation is also referred to as a hydrofinishing step. This step is suitably carried out at a temperature between 180 and 380 °C, a total pressure of between 10 to 250 bar and preferably above 100 bar and more preferably 35 between 120 and 250 bar. The WHSV (Weight hourly space

velocity) ranges from 0.3 to 2 kg of oil per litre of catalyst per hour (kg/l.h). Optionally the hydrogenation is performed in the same reactor as the catalytic dewaxing reactor. In such a reactor the beds of dewaxing catalyst and hydrogenation catalyst will be placed in a stacked bed on top of each other.

The hydrogenation catalyst is suitably a supported catalyst comprising a dispersed Group VIII metal. Possible Group VIII metals are cobalt, nickel, palladium and platinum. Cobalt and nickel containing catalysts may also comprise a Group VIB metal, suitably molybdenum and tungsten. Suitable carrier or support materials are low acidity amorphous refractory oxides. Examples of suitable amorphous refractory oxides include inorganic oxides, such as alumina, silica, titania, zirconia, boria, silica-alumina, fluorided alumina, fluorided silica-alumina and mixtures of two or more of these.

Examples of suitable hydrogenation catalysts are nickel-molybdenum containing catalyst such as KF-847 and KF-8010 (AKZO Nobel) M-8-24 and M-8-25 (BASF), and C-424, DN-190, HDS-3 and HDS-4 (Criterion); nickel-tungsten containing catalysts such as NI-4342 and NI-4352 (Engelhard) and C-454 (Criterion); cobalt-molybdenum containing catalysts such as KF-330 (AKZO-Nobel), HDS-22 (Criterion) and HPC-601 (Engelhard). Preferably platinum containing and more preferably platinum and palladium containing catalysts are used. Preferred supports for these palladium and/or platinum containing catalysts are amorphous silica-alumina. Examples of suitable silica-alumina carriers are disclosed in WO-A-9410263. A preferred catalyst comprises an alloy of palladium and platinum preferably supported on an amorphous silica-alumina carrier of which the commercially available catalyst C-624 of Criterion Catalyst Company (Houston, TX) is an example.

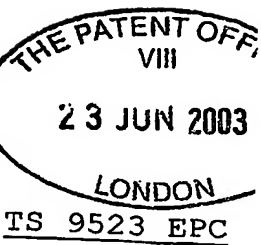
With the process according to the present invention different base oil grades may be prepared, such as spindle oil, light machine oil and medium machine oil having a saturates content of above 90 wt%, more preferably higher than 95 wt%. In the context of the present invention terms as spindle oil, light machine oil and medium machine oil will refer to base oil grades having an increasing kinematic viscosity at 100 °C and wherein the spindle oil additionally has a maximum volatility specification. Preferably a spindle oil is a light base oil product having a kinematic viscosity at 100 °C of below 5.5 cSt and preferably above 3.5. The spindle oil can have either a Noack volatility, as determined by the CEC L-40-T87 method, of preferably below 20% and more preferably below 18% or a flash point, as measured according to ASTM D93, of above 180 °C. Preferably the light machine oil has a kinematic viscosity at 100 °C of below 9 cSt and preferably above 6.5 cSt and more preferably between 8 and 9 cSt.

Preferably the medium machine oil has a kinematic viscosity at 100 °C of below 13 cSt and preferably above 10 cSt and more preferably between 11 and 12.5 cSt. The corresponding base oil grade can have a viscosity index of between 95 and 120.

The above referred to base oils are typically API Group II base oils having a viscosity index of between 80 and 120. With the present invention it is also possible to prepare so-called API Group III base oils having a viscosity index of above 120 by for example adding more of the Fischer-Tropsch derived fraction in step (b), adjusting the process conditions in step (a) or by using a crude derived feedstock which in itself yields a higher VI base oil. In the context of the present invention the content of the Fischer-Tropsch derived fraction in the

mixture obtained in step (b) is less than 60 wt%, preferably less than 50 wt%.

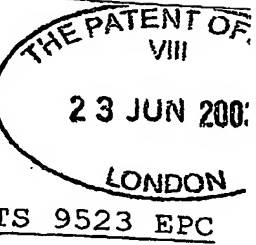
The above base oil grades may be obtained by distilling the product as obtained after step (c) or 5 step (d). In some base oil processing units comprising hydrocracking and catalytic dewaxing these base oil grades are prepared one at a time in a so-called blocked out mode as for example described by Figure 1.1 on page 2 of the above referred to General Textbook of Avilino 10 Sequeira Jr. Another option is that a full range feed is processed in step (a) and that from the effluent of step (a) fractions are isolated which correspond to the above spindle, light and medium machine oil grade as for example described in the above referred to WO-A-0250213. 15 The individual grades are subsequently further processed in step (c) in a blocked out mode. In terms of the present invention one or more of these grades can be mixed with the Fischer-Tropsch fraction. When processing the different grades separately through steps (a) and/or 20 (c) it is possible to use the Fischer-Tropsch fraction to correct only for those grades which need correction in VI. In a prior art process without having this possibility it was not possible to target the desired VI for every grade. In practice one would target the VI for 25 the most difficult grade and accept a VI much higher than the specification for the remaining grades. As explained above a too high VI implicates a non-optimal yield for the base oil. This quality give-away can now be avoided with the process according to the present process.



C L A I M S

1. Process to prepare a base oil having a viscosity index of above 80 and a saturates content of above 90 wt% from a crude derived feedstock by
  - 5 (a) contacting a crude derived feedstock in the presence of hydrogen with a catalyst comprising at least one Group VIB metal component and at least one non-noble Group VIII metal component supported on a refractory oxide carrier;
  - 10 (b) adding to the effluent of step (a) or part of the effluent of step (a) a Fischer-Tropsch derived fraction boiling at least partly in the base oil range in an amount effective to achieve the target viscosity index of the final base oil; and
  - 15 (c) dewaxing the mixture as obtained in step (b).
2. Process according to claim 1, wherein the crude derived feedstock is a vacuum distillate fraction or a de-asphalting vacuum residue as obtained from the residue of the atmospheric distillation of a crude petroleum feed.
- 20 3. Process according to any one of claims 1-2, wherein the viscosity index of the crude derived feedstock is below 60.
4. Process according to any one of claims 1-3, wherein the conversion in step (a) is between 20 and 80 wt%.
- 25 5. Process according to any one of claims 1-4, wherein in step (a) the crude derived feedstock is first subjected to a hydrotreating step prior to the hydrocracking step.
6. Process according to claim 5, wherein the conversion in the hydrotreating step is below 30 wt%.

7. Process according to any one of claims 1-6, wherein the kinematic viscosity at 100 °C of the mixture as obtained in step (b) is between 3 and 10 cSt.
8. Process according to any one of claims 1-7, wherein  
5 step (c) is performed by means of catalytic dewaxing.
9. Process according to any one of claims 1-8, wherein the dewaxed product of step (c) is subjected to an additional hydrogenation treatment step (d).
10. Process according to any one of claims 1-9, wherein the Fischer-Tropsch derived fraction is obtained by hydroisomerization of a Fischer-Tropsch synthesis product.



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### A B S T R A C T

#### PROCESS TO PREPARE A LUBRICATING BASE OIL

Process to prepare a base oil having a viscosity index of above 80 and a saturates content of above 90 wt% from a crude derived feedstock by

(a) contacting a crude derived feedstock in the presence of hydrogen with a catalyst comprising at least one Group VIB metal component and at least one non-noble Group VIII metal component supported on a refractory oxide carrier;

(b) adding to the effluent of step (a) or part of the effluent of step (a) a Fischer-Tropsch derived fraction boiling at least partly in the base oil range in an amount effective to achieve the target viscosity index of the final base oil; and

(c) dewaxing the mixture as obtained in step (b);